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- Novel vinyl-terminated carbamylmethylated melamines, vinyl-terminated polyurethane/polyamide polymers and coating/adhesive compositions containing same.

Disclosed are novel nitrogen containing heterocyclic compounds and novel vinyl terminated polyurethane/polyamide polymers together with the use of both of these compounds in a novel adhesive and coating composition. The nitrogen containing heterocyclic compounds of the invention contain two vinyl-terminated substituents and at least one carbamylmethyl substituent and have a nucleus selected from melamine, oligomers of melamine, benzoguanamine, and oligomers of benzoguanamine, glycoluril and oligomers of glycoluril. The vinyl terminated polyurethane compounds of the invention have a molecular weight of from about 3000 to about 80000, and contain at least two vinyl end groups.

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NOVEL VINYL-TERMINATED CARBAMYLMETHYLATED MELAMINES, VINYL-TERMINATED POLYURETHANE/POLYAMIDE POLYMERS AND COATING/ADHESIVE COMPOSITIONS CONTAINING SAME

The present invention relates to novel nitrogen containing heterocyclic compounds containing reactive functionalities inclusive of vinyl-terminated groups and carbamylmethyl groups. These compounds have general utility as monomers and crosslinking agents. This invention also includes novel vinyl-terminated polyurethane/polyamide polymers having a molecular weight of from about 3,000 to about 80,000 and containing at least two vinyl end groups. This invention also includes coating/adhesive compositions prepared from both (i) the novel nitrogen containing heterocyclic compounds of the invention, and (ii) the novel vinyl terminated polyurethane/polyamide polymers of the invention, said compositions having low temperature cure capacity.

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Background of the Invention

The prior art has described curable compositions containing melamine and benzoguanamine derivatives as crosslinking agents.

- U.S. Patents 4,708,984 and 4,710,542 describe respectively beta-hydroxyalkylcarbamylmethylated aminotriazines and alkylcarbamylmethylated aminotriazines containing curing compositions. These curing agents show generally superior detergent and salt spray resistant properties and are preferably cured at temperatures of from 150°C. to about 200°C.
- U.S. Patents 4,230,740 and 4,230,550 describe melamine compounds with pendant vinyl terminated groups and the use of such compounds in radiation cured coatings.
- U.S. Patent 4,295,909 describes adhesive compounds using a urethane acrylate capped prepolymer based on polybutadiene polyol or a polyamine.
- U.S. Patent 3,855,379 describes vinyl-terminated polyurethane polymers having a molecular weight of 2,000 to about 10,000 in bonding compositions.

Although the prior art describes melamine compounds with carbamylmethyl groups or melamine compounds with vinyl-terminated groups there is no disclosure of melamine compounds having both types of groups, or both types of groups in combination with methylol/alkoxymethyl groups on a melamine-type nucleus. Moreover, the prior art does not describe the combined use of vinyl-terminated polyurethanes/polyamides and melamine compounds with vinyl-terminated/carbamylmethyl groups in adhesive and coating compositions.

Summary of the Invention

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It has now been discovered that novel compounds having highly versatile and useful functionalities are nitrogen containing heterocyclic compounds having as essential pendant reactive groups; (i) at least 2 vinyl-terminated organic groups, and (ii) at least 1 carbamylmethyl group. Typically the nitrogen containing heterocyclic compounds containing the above groups are selected from melamine, oligomers of melamine, benzoguanamine, oligomers of benzoguanamine, glycoluril, oligomers of glycoluril, and mixtures thereof.

It is a further discovery of this invention that novel vinyl-terminated polyurethane/polyamide polymers having a molecular weight of from about 3,000 to about 80,000 containing at least two vinyl end groups have advantageous properties when used in combination with the nitrogen containing heterocyclic compounds of this invention.

It is a discovery of this invention that the novel heterocyclic nitrogen compounds and vinyl-terminated urethane compounds described in the preceding paragraphs are useful as essential ingredients in novel and improved low-temperature curing adhesive and coating formulations. This invention provides adhesives suitable for bonding, glass to glass, steel to steel, steel to glass, and plastic materials, including fiber reinforced sheet molding compound (SMC), to the same plastics, with little or no surface pretreatment.

DETAILED DESCRIPTION OF THE INVENTION

PART I - THE NOVEL NITROGEN CONTAINING HETEROCYCLIC COMPOUNDS:

The novel compounds of the invention are nitrogen containing heterocyclic compounds containing at least 2 vinyl-terminated substituents and at least 1 carbamylmethyl substituent. Also novel are compositions of matter which contain a predominant proportion of one or more novel compounds of the invention.

The nucleus of the nitrogen containing heterocyclic compound of the invention is desirably selected from melamine, oligomers of melamine, benzoguanamine, oligomers of benzoguanamine, glycoluril, and oligomers of glycoluril. A nucleus derived from melamine, oligomers of melamine, benzoguanamine, or oligomers of benzoguanamine is preferred. A nucleus derived from melamine or oligomers of melamine is most preferred. As used herein the word "oligomer(s)" of melamine, benzoguanamine, and glycoluril refers to amino resins prepared from these compounds by reaction with aldehydes such as formaldehyde.

The sites on the nitrogen containing heterocyclic compound nucleus not occupied by vinyl-terminated substituents or carbamylmethyl substituents may be occupied by any non-interfering substituents but it is preferred that such non-interfering substituents be methylol and/or alkylated methylol groups.

Novel Compounds based on Melamine or Melamine Oligomers:

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The novel melamine type compounds of the invention are represented by Formula (I) below: $C_3N_5(CH_2)_a(H)_{(6-a)}(R^1)_m(R^2)_n(R^3)_{a\cdot(n+m)};$ (I) where a=3 to 6; wherein R^1 is a substituent selected from the group

CH₂=C-CNH

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and

and

R4 CH=C-CH₂-O- , or mixtures thereof, and wherein Z is Ch2CH2-O- or

and wherein R4 is a hydrogen or a C1 to C18 alkyl radical, and wherein R2 is a carbamyl radical of the

Ö R5-O C-NH-

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wherein R5 is a C1 to C18 alkyl, alicyclic, hydroxyalkyl, alkoxyalkyl or aromatic radical, and wherein R3 -OR⁶, wherein R⁶ is a hydrogen or C₁ to C₁₈ aliphatic, alicyclic, or aromatic radical; with the proviso that m is at least 2, n is at least 1, and the sum of (m+n) is at least 3.

The preferred nitrogen containing heterocyclic compounds of the invention may alternatively be represented by Formula (II) below:

(II)

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wherein the substituent R has the following identity:

R is acrylamido, methacrylamido, acrylate, methacrylate, -OR5 where R6 is as defined previously, or carbamyl; provided that at least one R substituent is carbamyl, and that at least two R substituents are selected from acrylamido, methacrylamido, allyloxy, methallyloxy, acryloyloxy, acryloyloxy alkoxy, methacryloyloxyalkoxy and methacryloyloxy.

Illustrative melamine type compounds of this invention are as follows:

N2,N4,N6-tris(acrylamidomethyl)-N2,N4,N6-tris(2-ethylhexylcarbamylmethyl)melamine

N²,N⁴,N⁶-tris(2-ethylhexylcarbamylmethyl)-N²,N⁴,N⁶-tris(2-acryloyloxyethoxymethyl)melamine

 $N^2, N^4 - bis (methyl carbamylmethyl) - N^2, N^4, N^6 - tetrak is (2-acryloyloxyethoxymethyl) melamine a significant of the control of the$

N2,N4-bis(methylcarbamylmethyl)-N2,N4,N5,N5-tetrakis(2-methacryloyloxyethoxymethyl)melamine

N²,N⁴,N⁶-tris(propylcarbamylmethyl)-N²,N⁴,N⁶-tris(2-acryloyloxyethoxymethyl)melamine

N²,N⁴,bis(propylcarbarnylmethyl)-N²,N⁴,N⁶,N⁶-tetrakis(2-acryloyloxyethoxymethyl)melamine

 N^2,N^4,N^6 -tris(propylcarbamylmethyl)- N^2,N^4,N^6 -tris(2-methacryloyloxyethoxymethyl)melamine

N2,N4,bis(methylcarbamylmethyl)-N2,N4,N6,N6-tetrakis(2-methacryloyloxypropyloxymethyl)melamine

N2,N4,N6-tris(acrylamidomethyl)-N2,N4,N6-tris(methylcarbamylmethyl)melamine

N2,N4,N6-tris(methylcarbamylmethyl)-N2,N4,N5-tris(2-acryloloxyethoxymethyl)melamine

 $N^2, N^4, N^6 - tris(methylcarbamylmethyl) - N^2, N^4, N^6 - tris(2-methacryloloxyethoxymethyl) melamine$

 $N^2, N^4, N^6 - tris(methylcarbamylmethyl-N^2, N^4, N^6 - tris(2-acryloloxypropyloxymethyl) melamine a simple of the control of the control$

N²,N⁴,N⁶-tris(propylcarbamylmethyl)-N²,N⁴,N⁶-tris(2-methacryloloxypropyloxymethyl)melamine

 $N^2, N^4-bis (propylcarbamylmethyl-N^2, N^4, N^6-tetrak is (2-methacryloloxypropyloxymethyl) melamine and the state of t$

 N^2,N^4 ,-bis(propylcarbamylmethyl)- N^2,N^4 , N^5 - N^6 -tetrakis(2-acryloloxyethoxymethyl)melamine

N²,N⁴,N⁶-tris(propylcarbamylmethyl)-N²,N⁴,N⁶-tris-(acrylamidomethyl)melamine,

and mixtures thereof.

Formula (I) corresponds to the melamine type compounds of the invention when m is at least 2, n is at least 1 and the sum of (m+n) is at least 3.

The novel melamine compounds of the invention include oligomeric forms of melamine represented by Formula (III) below:

wherein the degree of polymerization, p is from 2 to 10, with values of p from 2 to 5 being preferred. Oligomers of melamine are commercially available as CYMEL® 300 series resins from American Cyanamid Company, Wayne, New Jersey. The melamine oligomers are typically joined by the linkage "L" shown above wherein L is a -CH₂- or -CH₂OCH₂-linkage which occupies sites on the melamine nucleus, however, the remaining reactive sites on the oligomeric nucleus must comply with the general requirement that at least 2 vinyl-terminated substituents and at least 1 carbamyl substituent be present.

Novel Compounds Based on Benzoguanamine and Benzoguanamine Oligomers:

The novel benzoguanamine type compounds of the invention are represented by Formula (IV) below: $(C_6H_5)C_3N_5-(CH_2)_b(H)_{4-b}-(R^1)_m(R^2)_n(H^3)_{b-(n+m)};$ (IV) where b=3 to 4; wherein R¹, R², R³, m and n are as defined in the preceding section.

Oligomeric forms of benzoguanamine having at least 2 vinyl and at least 1 carbamyl group are also compounds within the scope of this invention.

Novel Compounds based on Glycoluril and Glycoluril Oligomers:

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The novel glycoluril type compounds of the invention are represented by Formula (III) below: $C_4H_2N_4(O)_2$ - $(CH_2)_c(H)_{4-c}$ - $(R^1)_m(R^2)_n(R^3)_{c-(m+n)}$; where c=3 to 4; wherein R^1 , R^2 , R^3 , n, and m are as defined in the preceding section.

Oligomeric forms of glycoluril having at least 2 vinyl-substituted and at least 1 carbamyl group are compounds within the scope of this invention.

Common Aspects of the Novel Nitrogen Containing Heterocyclic Compounds of the Invention:

The vinyl groups have been described in general terms, but may also be referred to as substituents derived from unsaturated alcohols, acids, and amides. Examples of compounds suitable for the second step of the vinyl-terminated oligomer preparation are allyl alcohol, hydroxy ethylhexyl acrylate, 2-hydroxyethyl methacrylate, 2-hydroxy propyl methacrylate, hydroxylauryl methacrylate, 2-hydroxyethyl acrylate, acrylic acid, methacrylic acid, ethacrylic acid, crotonic acid, methacrylamide, N-hydroxy acrylamide, N-hydroxy methacrylamide, acrylamide, and mixtures thereof.

The carbamylmethyl groups have been described in general terms, but may also be referred to as substituents derived from alkyl and hydroxy alkyl carbamates of from 1 to 18 carbon atoms. Examples of suitable carbamate reactants used to produce the carbamylmethyl substituents of the invention are as follows: methyl carbamate, ethyl carbamate, propyl carbamate, butyl carbamate, beta-hydroxy alkyl carbamates, 2-ethylhexyl carbamate, and beta-alkoxy alkyl carbamates.

The minimum number of vinyl-terminated substituents and carbamylmethyl substituents has been stated to be 2 and 1, respectively. However, it is generally advantageous to have a plurality of both vinyl-terminated and carbamylmethyl substituents on the compounds of the invention. In the case of melamine systems, it is desirable that each N-containing heterocyclic ring contains at least three or more vinyl groups and one or more carbamyl groups. Preferably, the heterocyclic ring should contain 3 to 5 vinyl groups and 1 to 3 carbamyl groups.

The Method of Making the Novel Nitrogen Containing Heterocyclic Compounds of the Invention:

The novel compounds of the invention may be prepared by the reaction of (A) alkoxymethylated or

methylolated nitrogen containing heterocyclic compounds with (B) reactants consisting (i) an alkyl carbamate, and (ii) a terminally unsaturated acid, alcohol, or amide.

The nitrogen containing heterocyclic ingredient (A) may be combined with reactants (B) (i) and (B) (ii) simultaneously or in any order. Thus, reaction with alkyl carbamate may first be effected and thereafter reaction with the vinyl-terminated compound, or vice versa. In accordance with the present invention, the starting materials are reacted with alkyl carbamates, such as methyl carbamate or propyl carbamate in the presence of an acid catalyst. An illustrative reaction is depicted by the following equation (for melamine type compounds):

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wherein vinyl is as defined above and wherein R⁵ and R⁶ are as previously defined and m is the desired degree of carbamylmethylation. Reaction is typically carried out by heating in the melt or in solution, e.g., in toluene, in the presence of catalytic amounts of acid, sulfuric acid, and the like at temperatures between 80°C. and 150°C. Measurement of the quantity of alcohol and/or water evolved gives an indication of the degree of reaction completion. Typically, for the carbamylmethylation of melamine m is from 2 to 4, and for the carbamylmethylation of benzoguanamine or glycoluril m is 2. The method of placing carbamylmethyl substituents on the nitrogen containing heterocyclic nucleus is generally described at columns 3 and 4 of U.S. Patent No. 4,710,542; the disclosure of which is incorporated herein by reference.

The method of placing vinyl-terminal groups on the nitrogen containing heterocyclic compounds may be found in Helv Chim Acta 24,318 E, (1941), A. Gams, G. Widmer and W. Fisch which shows a transetherification method of making allylic terminated types and U.S. Patent No. 3,020,255 which teaches a transetherification method of making acrylic terminated derivatives. In carrying out the transetherification reaction the degree of substitution n of the vinyl-terminated groups is adjusted to be typically from 2 to 4 in the case of melamine and n equals 2 to 3 in the case of benzoguanamine or glycoluril. Suitable vinyl reactants are selected from acrylamide, methacrylic acid, methacrylamide, acrylic acid, and hydroxy functional monomers such as hydroxy alkyl acrylates and hydroxy alkyl methacrylates. The disclosure of U.S. Patent 3,020,255 at columns 4 to 5 is incorporated herein by reference.

The reactions required to place vinyl-terminated groups and carbamyl groups onto the alkylmethylolated methylolated heterocyclic nucleus may be carried out simultaneously or separately and may be performed in any desired sequence.

The method of making the novel nitrogen containing heterocyclic compounds of this invention as described in this section yields a mixture of products which represent a statistical distribution of vinyl-terminated substituents and carbamylmethyl substituents. Typically, at least the predominant part by weight of the reactor product contains novel nitrogen containing heterocyclic compounds having at least two vinyl-terminated substituents and at least one carbamylmethyl substituent.

The novel compounds of the invention have general utility as resin precursors for the formation of objects and coatings, however, it is a discovery of this invention that the above described nitrogen containing heterocyclic compounds containing both vinyl-terminated substituents and carbamylmethyl substituents have special utility in preparing novel adhesive, coatings and sealant compositions. Moreover, a composition of matter which is a mixture of the novel nitrogen containing heterocyclic compounds of the invention (e.g., as prepared by the methods of this section) has special utility in preparing novel coatings, adhesives, and sealants.

PART II - THE NOVEL VINYL-TERMINATED POLYURETHANE/POLYAMIDE POLYMERS

The adhesive composition of the invention contains from about 30 to 90 weight percent of a vinyl-terminated polyurethane having a molecular weight of from about 3,000 to about 80,000, preferably 3,000 to 7,000, containing at least two vinyl end groups.

The novel vinyl-terminated polyurethane/polyamide component of the invention may be prepared by first reacting a polyisocyanate with a vinyl acid, vinyl alcohol, or vinylamide, then as a second step reacting the first step reaction product with a high molecular weight polyol.

The isocyanate material used to produce the isocyanate terminated polyurethane resin includes aliphatic, cycloaliphatic, and aromatic diisocyanates such as 2,4-tolylene diisocyanate, metaphenylene diisocyanate, 2,6-tolylene diisocyanates, p.p'-diphenylmethane diisocyanate, 1,5-naphthylene diisocyanate, phenyl diisocyanate, dianisidine diisocyanate, 1,5-naphthalene diisocyanate, 4,4'-diphenyl ether diisocyanate, p-phenylene diisocyanate, 4,4'-dicyclo-hexylmethane diisocyanate, 1,3-bis(isocyanatomethyl)-cyclohexane, cyclohexylene diisocyanate, tetrachlorophenylene diisocyanate, isophorone diisocyanate, 2,6-diethyl-p-phenylenediisocyanate, 3,5-diethyl-4,4'-diisocyanatodiphenyl-methane, toluene diisocyanato-4,4'-diphenyl diisocyanate, phenylene diisocyanates, octamethylene diisocyanate, 3,3'-dimethoxy-4,4'-biphenyl diisocyanate, hexamethylene diisocyanate, tetramethyl xylylene diisocyanate (viz., TMXDI® polyisocyanate available from American Cyanamid Company, Wayne, New Jersey). The preferred isocyanates are tetramethyl xylylene diisocyanate, 2,4-or 2,6-tolylene diisocyanates (2,4-TDI or 2,6-TDI), 4,4'-diphenyl-methane diisocyanate (MDI)., isophorone diisocyanate (IPDI), and hexamethylene diisocyanate (HMDI).

Compounds suitable for effecting the first step reaction include vinyl alcohols, vinylic acids, and vinyl amides, hydroxy acrylates, hydroxy alkyl acrylates, and mixtures thereof. Examples of compounds suitable for the first step of the vinyl-terminated oligomer preparation are allyl alcohol, hydroxy ethylhexyl acrylate, 2-hydroxyethyl acrylate, propyl methacrylate, 2-hydroxypropyl acrylate, 2-hydroxybutyl acrylate, w-hydroxybutylacrylate, p-hydroxyphenyl acrylate, hydroxylauryl methacrylate, 2-hydroxypropyl methacrylate, 3-hydroxypropyl methacrylate, hydroxyhexyl acrylate, hydroxylauryl methacrylate, the monoacrylate or monmethacrylate esters of bisphenol A, the fully hydrogenated derivatives of bisphenol A, polyethyleneglycol methacrylate, acrylic acid, methacrylic acid, ethylacrylic acid, crotonic acid, methacrylamide, acrylamide, and mixtures thereof.

The second step high molecular weight polyol reactant is typically selected from polyhydroxy functional polymeric materials which include but are not limited to polycaprolactone diols, hydroxy terminated polybutadiene polymers, hydroxy terminated poly(butadiene-acrylonitrile) polymers, hydroxy terminated poly(butadiene-styrene) polymers, multitudes of polyester polyols and polyether polyols are also suitable for this invention. The selection of the optimum polyol is dictated by specific end-user's applications. For example, the preferred polyols for plastic substrates (e.g. SMC) are the hydroxy-terminated polybutadiene and poly(butadiene-acrylonitrile) copolymers.

It is permissible to reverse the order of reaction used in the preparation of the vinyl-terminated polyurethane/polyamide polymer. Thus, in a first step reaction polyisocyanate compound is reacted with a high molecular weight polyol. Thereafter, the first step reaction product is reacted with a vinyl acid, vinyl alcohol, or vinyl amide to prepare a vinyl-terminated polyurethane/polyamide polymer substantially free of unreacted isocyanate substituents. The final vinyl-terminated polyurethane may also have additional vinyl substitution at internal positions in its structure, if desired.

The mole ratio of isocyanate groups to active hydrogen groups (e.g., $-NH_2$; -OH; -COOH) in the first step reaction is from about 1:1 to 1.1:1 with ratios of isocyanate groups to active hydrogen groups in the first plus second steps of 1:1 being preferred.

Generally, each of the two reaction steps are carried out at temperatures from 50°C. to 125°C. for a time-between 0.5 to 4 hours. The preferred conditions are in the range of 60-95°C. for 2 to 3 hours. The process may be operated in either a batch or continuous mode.

PART III - THE NOVEL COATING, SEALANT, AND ADHESIVE COMPOSITIONS:

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The composition of the invention have general utility as coatings, sealants, and adhesives. All compositions of this invention contain the following essential ingredients:

- (a) at least one nitrogen containing heterocyclic compound having as reactive groups; at least 2 vinyl terminated organic groups, and at least one carbamylmethyl group.
- (b) at least one vinyl-terminated polyurethane/polyamide polymer having a molecular weight of from about 3,000 to about 80,000 as described in PART II of this specification.

The coating compositions and sealant compositions of this invention have as essential ingredients components A, B, C, and D as follows:

- A. from about 5 to about 70 weight percent of the vinyl-terminated, cabamylmethylated nitrogen containing heterocyclic compounds of the invention.
 - B. from about 30 to about 90 weight percent of a vinyl-terminated polyurethane/polyamide polymer

having a molecular weight of from about 3,000 to about 80.000;

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- C. from about 2 to about 30 weight percent of a polymerizable diluent;
- D. from about 0.5 to about 5 percent of a free-radical catalyst, wherein the sum of ingredients A, B, C and D does not exceed 100 weight percent. Preferably, the weight percent limits of the essential coating/sealant composition ingredients are as follows:
- A. from 15 to 40 weight percent of the vinyl-terminated, carbamylmethylated nitrogen containing heterocyclic compound.
 - B. from 50 to 70 weight percent of the vinyl-terminated polyurethane/polyamide polymer.
 - C. from 5 to 15 weight percent of the polymerizable diluent;
 - D. from 0.5 to 3.0 weight percent of the free-radical catalyst.

The adhesive compositions fo this invention have as essential ingredients components A, B, C, D, and E as follows:

- A. from about 5 to about 70 weight percent of the vinyl-terminated, carbamylmethylated nitrogen containing heterocyclic compounds of the invention.
- B. from about 30 to about 90 weight percent of a vinyl-terminated polyurethane/polyamide polymer having a molecular weight of from about 30,000 to about 80,000;
 - C. from about 2 to about 30 weight percent of a polymerizable diluent;
 - D. from about 0.5 to about 5 weight percent of a free-radical catalyst;
 - E. from about 0.5 to about 10 weight percent of an adhesion promoter/coupling agent, wherein the sum of ingredients A, B, C, D and E does not exceed 100 weight percent. Preferably, the weight percent limits of the essential adhesive composition ingredients A, B, C, and D are the same as those set out for the coating/sealant composition previously described. The preferred content of ingredient E, the adhesion promoter, is from 2 to 7 weight percent.

Ingredient (A) - the nitrogen containing heterocyclic compound:

The vinyl-terminated carbamylmethylated nitrogen containing heterocyclic compounds of the invention are those described at "Part I - THE NOVEL NITROGEN CONTAINING HETEROCYCLIC COMPOUNDS:" of this disclosure. These novel compounds may be used separately or in any combination as ingredient (A) of adhesive composition described herein. In particular, ingredient (A) may be a composition of matter containing a predominant part by weight of a statistical distribution of novel nitrogen containing heterocyclic compounds of the invention, wherein said compounds have at least two vinyl-terminated substituents and at least one carbamylmethyl substituent. The novel compounds based on melamine and oligomers of melamine are particularly preferred for the formulation of compositions according to this invention.

Ingredient (B) - the novel vinyl-terminated polyurethanes/polyamides:

The vinyl-terminated polyurethane/polyamides ingredient of the invention is described at "Part II -THE NOVEL VINYL-TERMINATED POLYURETHANE/POLYAMIDES POLYMERS" of this disclosure. These novel compounds are used as ingredient (B) of compositions described herein.

5 Ingredient (C) - the Polymerizable Diluent:

The composition of the invention includes a polymerizable diluent that does not require removal during use or cure. The diluent participates in the formation of coating, sealant, or adhesive and is substantially incorporated into the cured composition.

Examples of reactive diluents useful in the practice of this invention are ethylene glycol diacrylate, ethylene glycol dimethacrylate, diethylene glycol diacrylate, diethylene glycol dimethacrylate, trimethylol propane triacrylate, trimethylol propane trimethacrylate, pentaerythritol triacrylate, pentaerythritol triacrylate, pentaerythritol triacrylate, neopentyl glycol diacrylate, isobutoxy methacrylate (IBMA), styrene, vinyl pyrrollidone, alpha methylstyrene, meta-diisopropenylbenzene, para-diisopropenylbenzene; and mixtures thereof. Particularly preferred as reactive diluents are styrene and vinyl pyrrollidone.

The proportion of polymerizable diluent is not critical although it usually constitutes from 2 to 30 weight percent and preferably 5 to 15 weight percent of the adhesive composition. The proportion of polymerizable

diluent may be adjusted to achieve a desired viscosity in the adhesive composition.

The polymerizable diluent is desirably also a solvent for the other ingredients of the adhesive composition. The term, solvent, as used herein means that the other ingredients are solubilized under the conditions of use of the composition.

Ingredient (D) - the Free-Radical Catalyst:

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The catalysts useful in the practice of the invention are selected from conventional free radical catalysts which decompose on heating. Generally suitable catalysts are those of the peroxide or azo type. Illustrative free-radical generating agents of the peroxide type include but are not limited to benzoyl peroxide, 2,4-dichlorobenzoyl peroxide, lauroyl peroxide, octanoyl peroxide, acetyl peroxide, dialkylperoxides such as ditert-butyl peroxide, dicumyl peroxide, ketone peroxides such as methylethylketone peroxide; and peresters which readily decompose, e.g., tertiary-butyl peracetate, tertiary butyl perbenzoate, ditertiary butyl diperphthalate, t-butyl peroxyisobutyrate, 2,5-dimethyl-2,5-di(benzoyl peroxy)hexane, 2,5-dimethyl-2,5-bis(tertiary butylperoxy)hexane-3, n-butyl-4,4-bis(tertiary butylperoxy) valerate. Another useful class of catalysts are peroxy compounds such as the organic hydroperoxides, for example, cumene hydroperoxide, cyclohexanone peroxide, methylethyl ketone hydroperoxide, tertiary butyl hydroperoxide.

The catalyst systems can optionally contain accelerations/promoters such as diethylaniline, dimethylaniline, condensation products of aldehydes and primary or secondary amines; metal salts of octyl acid, stearic acid, oleic acid, linoleic acid, naphthenic acid and rosin acid and said metal is selected from chromium, iron, cobalt, nickel, manganese and lead, lauroyl mercaptan, thiourea. Typically, from 0.1 to 5 weight percent of accelerator/promoter is used.

Particularly preferred catalysts of this invention are the azo type free radical catalysts such as alphaalpha azo-di-isobutylronitrile.

Preferred azo polymerization initiators have the following molecular structures:

methyl, ethyl, or isobutyl; and

$$H_{2}C$$

$$CH_{2}-CH_{2}$$

$$CH_{2}-CH_{2}$$

$$CH_{2}-CH_{2}$$

$$CH_{2}-CH_{2}$$

$$CH_{2}-CH_{2}$$

$$CH_{2}-CH_{2}$$

Typically, the free-radical catalysts constitute from about 0.5 to about 10.0 weight percent of the composition. The activity of the catalyst as measured by its half-life is a useful indicator of the suitability for the free radical generator for a particular application. The utility of a particular free radical catalyst may be judged by mixing it with the other components of the composition of the invention and determining the resultant shelf-life and degree of cure at the temperature of use. The catalyst may be used in microencapsulated form if desired.

Ingredient (E) - the adhesion promoter/coupling agent:

The adhesive composition of the invention contains ingredients for enhancing its bonding to substrates such as steel, glass and plastics.

Useful adhesion promoting agents include silane compounds and organic anhydrides.

Silane coupling agents improve the moisture resistance, adhesion, and durability of the adhesive or

coating composition prepared herein. As silane coupling agents for the present invention there may be used any conventional silane coupling agent. Preferred are silane coupling agents having a vinyl group since such materials can participate in the polymerization reactions which cure the composition of the invention. The silane compounds are typically used at concentrations of from about 0.5 to about 10.0 weight percent of the composition. Silane compounds of the following generic formula are preferred. X-CH₂CH₂CH₂Si(OR)₃

wherein X is NH_2 , R is C_2H_5 , or X is $CH_2 = C(CH_3)CO_2$, and R is CH_3 .

Examples of suitable silane type coupling agents are methacryloxypropyltrimethoxysilane, vinyltriethoxy silane, vinyltrimethoxy silane, vinyltris(beta-methoxyethoxy)silane, and mixtures thereof.

Stoichiometric amounts of water relative to hydrolyzable alkoxysilane groups should be present if silane compounds are incorporated into the composition without pretreatment or substrate surfaces.

The adhesion promoter may be an anhydride such as succinic anhydride, maleic anhydride, alkyl/alkenyl succinic anhydride, e.g., dodecenyl succinic anhydride, tetrahydrophthalic anhydride, hexahydrophthalic anhydride, phthalic anhydride, glutaric anhydride, 1,4,5,6,7,7-hexachlorobicyclo[2.2.1]-5-heptene-2,3-dicarboxylic anhydride, tetrachlorophthalic anhydride, 3,3,4,4,-benzophenone tetracarboxylic dianhydride, polyazelaic polyanhydride, pyromellitic dianhydride, pyromellitic dianhydride-glycol adducts, 1,2,3-cyclopentanetertcarboxylic dianhydride and mixtures thereof. Anhydrides are especially useful adhesion promoters when used with SMC.

Optional Ingredients:

Additives such as stabilizers, accelerators, activators, flame retardants, pigments, plasticizers, surface active agents, dyestuff, inert filler, and other ingredients which do not adversely affect cure may be incorporated into the composition.

Packaging of Ingredients:

The composition of the invention has the advantage that it may be formulated as a one-package system that has acceptable shelf-life and may be used "as is" at the time coating or adhesion is required by the user. Typically, a one-package system comprises a mixture of the essential ingredients (A, B, C, D for coating compositions or A, B, C, D, E for adhesive compositions) as described in this section.

The mixture of essential ingredients of the compositions of the invention is typically and advantageously in the form of a solution wherein the ingredients are mutually soluble under conditions of use. It is possible, however, to use the curable compositions of this invention in the form of a suspension provided that the ingredients are uniformly dispersed with one another under conditions of use.

The one package system may also contain a stabilizer to delay or prevent crosslinking of the composition ingredients during transport or storage. The stabilizers useful for this invention include hydroquinones, p-benzoquinones, anthraquinones, naphthaquinones, phenanthraquinones and substituted compound of any of the foregoing. Additionally, various phenols can be employed such as 1,6-di-tert-butyl-4-methyl phenol, 4-methoxy phenol, 3,5-di-tert-butyl-hydroxytoluene.

The compositions of the invention may, however, be formulated as system two component packages. Typically, such a two-package system will comprise a first package containing the free-radical initiator, and the second package containing an accelerator/promoter for the selected free-radical catalyst. The two-package adhesive formulation is used by combining the packages at a time proximate to use.

Method of Use:

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The adhesive, sealant, and coating compositions of the invention are applied by any conventional method such as brushing, spraying, dipping, padding, or injection. The compositions are cured by application of heat, typically by convection ovens, microwave heating, or infra-red radiation. Generally, with conventional free-radical catalysts cure is effected at a temperature of from about 60 °C. to 150 °C.. In certain instances for bonding transparent substrates (e.g., glass, certain plastics), the curing may be effected by radiation, such as ultraviolet light or ultraviolet light in combination with heat. In general surface coating applications, the curing may be effected by radiation with incorporation of radiation-sensitive catalysts. Such radiation-sensitive catalysts include camphorquinone (1,7,7-trimethyl-2,3-diketo dicyclo-

(2,2,1-heptane)), benzil dimethyl ketal (BDMK) benzophenone (BP), diethoxyacetophenone (DEAP), hydroxycyclohexyl phenyl ketone (HCPK), benzil diethyl ketal(2,2-diethoxy-2-phenyl acetophenone), 2-isopropylthioxanthone, 2-chlorothioxanthone,1-phenyl-1,2-propanedione-2-(o-ethoxycarbonyl)oxime, di-sec-butoxy acetophenone, benzoin isopropyl ether, methyl phenyl glyoxylate, 2-hydroxy-2,2-di-methyl acetophenone, benzoin isobutyl ether.

The preferred method of curing the compositions of the invention is by application of heat. The time of cure will depend on the specific catalyst used, but generally it is desirable to have partial cure develop within one minute.

A particular advantage of the adhesive composition of the invention is that it may be applied to the substrates to be bonded without special surface preparation. Typically, a methylene chloride wipe or light abrasion may be used as standard surface preparation. In many instances the adhesive compositions of this invention may be applied to soiled surfaces and give acceptable results.

Objects formed by the Compositon of the Invention:

The composition of the invention is useful in forming composites, fiber-reinforced composites laminates, adhesive interlayers, sealants, coatings, films, and solid objects. The most frequent application will be for the formation of adhesive interlayers and topcoatings.

The invention is illustrated by the following Examples.

PART ONE - PREPARATION OF THE NOVEL HETEROCYCLIC NITROGEN COMPOUNDS OF THE INVENTION,

Ingredient (A)

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EXAMPLE 1

This Example illustrates the preparation of the novel compounds of the invention.

A 2 liter flask was flushed with dry air and charged with the following ingredients:

158.4 grams of hexamethoxymethylmelamine (CYMEL® 300, product of American Cyanamid Company, Wayne, New Jersey)

210.9 grams of 2-ethylhexyl carbamate

86.4 grams of acrylamide

1.35 grams of hydroquinone

600 milliliters of cyclohexane

The reaction flask was equipped with a condenser and trap. Initially the flask contents were heated in a 90°C. bath until they became clear, then 4.5 grams of para-toluenesulfonic acid was added to the flask.

The reaction was monitored by watching the methanol byproduct accumulate in the trap. When methanol collection stopped, methanol was removed from the trap and weighed.

The reaction was concluded when the amount of methanol collected reached the approximate theoretical methanol yield. For this reaction the theoretical methanol yield was achieved about three and one-half hours after the addition of the para-toluenesulfonic acid.

The cyclohexane was removed by rotary evaporator and the flask contents reconstituted with 500 milliliters of CH2 Cl2. Thereafter, the para-toluenesulfonic was neutralized with 100 milliliters of 5% by weight sodium bicarbonate solution. The organic layer was separated and dried over anhydrous potassium monocarbonate. Finally, the product was filtered and the solvent removed under vacuum (30 °C. at 10 millimeters Hg pressure) to give a final product of melamine acrylamide carbamate (hereinafter called compound "M".)

The product yield was 350 grams mainly consisting of melamine substituted with 3 carbamyl-methyl substitutents and 3 acrylamide substituents (tris-2-ethylhexyl-carbamylmethyl, tris-acrylamido-methyl melamine).

EXAMPLE 2

A 250 ml flask was flushed with dry air and charged 19.5g of Cymel® 300, 21.6g of 2-ethylhexyl 5 carbamate, 0.012g of phenothiazine and heated to 95 °C. for 15 minutes to get a clear solution. Thereafter, 0.28g of p-toluenesulfonic acid was added and heating was continued for 15 minutes. The pressure was gradually lowered to 50 mm. Hg to collect MeOH distillate. When the theoretical amount of distillate was collected, the reaction mixture was cooled to 65°C. and 15.0g of 2-hydroxyethyl acrylate was added to the reactor with stirring. The pressure was reduced to 50 mm. Hg to collect MeOH byproduct. At the end of 45 minutes, no more MeOH came out and the reaction was stopped. To the cooled reaction products was added 100 ml of CH2Cl2. Two separate 25 ml washes of 5% wt. Na2CO3 solution was used to neutralize ptoluenesulfonic acid. The organic layers were separated and dried over anhydrous K2CO3. The above mixture was filtered and the solvent was removed to give 42g of a final product of melamine acrylate carbamate (hereinafter called compound "M1").

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EXAMPLE 3

To a 250 ml 3-neck round-bottomed flask was charged 39g of Cymel® 300, 0.018g of phenothiazine and 30g of 2-hydroxyethylacrylate. A distillation head and mechanical stirrer were connected to the flask. The system was flushed with dry air.

The contents of the reaction flask were heated at 65°C. and 0.35g of p-toluenesulfonic acid (p-TSA) immediately added thereto. The reaction was run at 65° C. under vacuum pressure of 30 mm. Hg to remove methanol byproduct. At the end of 45 minutes, 8.6g of methanol was collected and the reaction stopped. To the cooled reaction product was added 50 ml of CH2Cl2, and the product was then washed with two 25 ml portions of 5% wt Na₂CO₃ solution. The CH₂Cl₂ solution of product was dried over anhydrous K₂CO₃. Filtration of solids and removal of solvent gave 48g products of melamine acrylates (hereinafter called compound "M2"). The product mainly consisted of melamine substituted with 3 ethylacrylate substituents and no carbamate groups.

EXAMPLE 4

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This Example illustrates the preparation of the novel compounds of the invention.

A 3 liter flask was flushed with dry air and charged with the following ingredients: 823.5 grams of hexamethoxymethylmelamine (CYMEL® 300, product of American Cyanamid Company,

316.5 grams of methyl carbamate

0.36 grams of phenothiazine

Wayne, New Jersey)

The reaction flask was equipped with a condenser and trap. Initially the flask contents were heated in a 75°C. bath for 15 minutes to get a clear solution, then 10.6 grams of para-toluenesulfonic acid was added to the flask.

The pressure of the reaction was gradually lowered from 200 to 50 millimeters of Hg to assist collection of metahnol distillate. After the theoretical amount of methanol distillate was collected (after about 60 minutes) the immersion oil bath around the flask was cooled to 65°C. Thereafter, 979 grams of 2hydroxyethyl acrylate was added with stirring to the reaction mixture. The pressure was again reduced gradually from 200 to 25 mm. of Hg and the methanol byproduct collected.

The reaction flask was cooled and 1200 milliliters of CH₂Cl₂ added to the contents. 20 grams of sodium bicarbonate dissolved in 300 milliliters of water was added to the flask contents to neutralize the remaining para-toluenesulfonic acid. The resultant organic reaction layer was separated and dried over anhydrous potassium carbonate.

The dried product was filtered and solvent removed by vacuum distillation at a pressure of 1 millimeter Hg at 30°C. to 40°C. to give a final product of melamine acrylate carbamate (hereinafter called compound "M3".)

The product yield was 1851 grams consisting of mainly melamine substituted with two carbamylmethyl substituents and four ethylacrylate substituents (bis-carbamylmethyl, tetrakis(2-acryloyloxyethoxy)-methyl melamine).

EXAMPLE 5

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A one liter flask was flushed with dry air and charged 205.8g of Cymel® 300, 295g of 2-hydroxypropyl-methacrylate (HPMA, Rocryl 410 from Rohm & Haas Company) and 0.1g of phenothiazine and heated to 75°C. for 15 minutes to get a clear solution. 2.6g of p-toluenesulfonic acid was added and heating continued for 15 minutes. The pressure was lowered to 50 mm. Hg for MeOH byproduct collection. When the theoretical amount of distillates was collected, the reaction was cooled to 65°C. 79g of methyl carbamate was charged and the reaction mixture stirred at 75°C. at 50 mm. Hg vacuum to collect MeOH. After 90 minutes, no more MeOH was generated and the reaction was stopped. To the cooled reaction products were added 300 ml of CH₂Cl₂ A two part 50 ml wash of 5% wt Na₂CO₃ solution was used to neutralize the p-TSA. The organic layers were separated and dried over anhydrous K₂CO₃. Filtration and removal of solvent gave 370g of final product of melamine methacrylate carbamate (hereinafter called compound "M4").

Other examples of resins containing carbamylmethyl and vinyl groups based on alkylated formaldehyde derivatives of melamine, benzogunamine and glycoluril are given in Examples 6 to 17. These Example preparations are based on the generalized procedure described below.

General Procedure

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A round bottom reaction flask was equipped with a mechanical stirrer and a trap with condenser suitable for vacuum distillation was charged with the reactants. The reaction mixture was heated in an 80°C. oil bath until the reactants were molten, then acid catalyst was added (usually 1-2 wt % H₂SO₄).

The reaction was monitored by the collection of alcohol via continuous vacuum distillation. The reaction was concluded when alcohol distillation had ceased. In cases where the amount of alcohol collected was below the theoretical, it was advisable to use a second addition of catalyst.

Because the reactivity fo carboxylic acids is low compared with carbamates and alcohols, it was desirable that a 20% to 40% excess of carboxylic acid be used. Any unreacted material was removable by increasing the vacuum before concluding the reaction.

After the contents of the reaction flask were cooled to near room temperature, 350 ml of CH₂Cl₂ was added in preparation for the work-up of the product. The methylene chloride solution was gently washed in a separatory funnel with 5% sodium bicarbonate solution to neutralize and extract the acid. The organic phase was separated and dried over anhydrous sodium sulfate until the solution had-cleared. Finally, the solution was vacuum filtered through the Buchner funnel and the solvent removed on a rotary evaporator.

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		TABLE OF TEST RESULTS FO	OR EXAMPLES 6 TO	10
1		CYMEL® HEXAMETHOXYMET	HYL MELAMINE SER	IES
t				Conditions
	Example	Reactants	Product substituents on Cymel/mole	temperature; time, catalyst
,	6	Cymel®-300, 150 g (0.385 mole) methyl carbamate, 57.8g (0.769 mole) acrylic acid, 110.9 g (1.54 mole)		75˚ C., 5 hrs. 0.8% H₂SO₄
	7	Cymel®-300, 150 g (0.385 mole) methyl carbamate, 57.8 g (0.769 mole) alpha-methyl acrylic acid, 132.5 g (1.54 mole)	145 g 2 carbamylmethyl 2 methacrylate	75 ° C.5.5 hrs. 1.0%H₂SO₄
5	8	Cymel®-300, 150 g (0.385 mole) n-propylcarbamate, 119.0g (1.16 mole) acrylic acid, 83.2 g (1.16 mole)	184 g 2.7 carbamyl propyl 1.9 acrylate	80 ° C.,4.0 hrs. 0.8% H₂SO₄
0	9	Cymel®-300, 150 g (0.385 mole) n-propylcarbamate, 119.0 g (1.16 mole) alpha-methyl acrylic acid, 99.3 g (1.16 mole)	175 g 1.7 carbamyl propyl 2.0 methacrylate	75° C., 8.0 hrs. 0.8%H₂SO₄
25	10	Cymel®-300, 150 g (0.385 mole) methyl carbamate, 578g (0.769 mole) acrylic acid, 166.4 g (2.31 mole)	190 g 2.0 carbamyl methyl 3.1 acrylate	70° C., 7.0 hrs. 0.8% H ₂ SO ₄ (grad. incr. to 1.35% H ₂ SO.

0		TABLE OF TEST RESULTS FO	R EXAMPLES 11 TO	14
		CYMEL® BENZOGUAN	AMINE SERIES	
;				Conditions
35	Example	Reactants	Product substituents on Cymel/mole	temperature; time, catalyst
	11	Cymel®-1123, 150 g (0.384 mole) methyl carbamate, 68.9 g (0.917 mole) acrylic acid, 66.2 g (0.917 mole)	144 g 2.0 carbamyl methyl 1.7 acrylate	75 [°] C., 5.5 hrs. 0.5% H₂SQ₄
40	12	Cymel®-1123, 150 g (0.384 mole) n-propyl carbamate, 47.3g, (0.459 mole) alpha-methyl acrylic acid, 120 g, (1.30 mole)	172 g 1.0 carbamyi propyi 2.8 methacrylate	80 ° C., 7.5 hrs. 0.8% H ₂ SO ₄ (grad. incr. to 2.0% H ₂ SO ₄)
45	13	Cymel®-1123, 100 g, (0.256 mole) 2-hydroxyethyl acrylate 106.5 g (0.917 mole) methyl carbmate 23.0 g (0.306 mole)	118 g 2.7 oxyethylacrylate 1.0 carbamyl methyl	80° C., 4.5 hrs. 1.0% H ₂ SO ₄
	14	Cymel®-1123m 100 g (0.256 mole) methyl carbamate, 19.2 (0.256 mole) acrylic acid, 100 g (1.30 mole)	102 g 1.0 carbamyi methyl 2.8 acrylate	80° C., 9 hrs. 1% H ₂ SO ₄ ; incr. to 1.7% H ₂ SO ₄

	CYMEL® 300 BUTYLATED GLYC	OLURIL SERIES	
			Conditions
Example	Reactants	Product substituents on Cymel/mole	temperature; time, catalyst
15	Cymel®-1170, 150g, (0.309 mole) methyl carbamate, 23.2 g (0.309 mole) alpha-methylacrylic acid, 93.0 g (1.08 mole)	144 g 1.0 carbamyl methyl 2.0 methacrylate	70° C., 5 hrs. 0.5% p-TSA
16	Cymel®-1170, 100g, (0.206 mole) methyl carbamate, 15.5 g (0.206 mole) hydroxyethylacrylate, 71.6 g, (0.617 mole)	111 g 1.0 carbamyl methyl 2.0 oxyethylacrylate	70° C., 4 hrs. 1.0% p-TSA 0.25% H₂SO₄
17	Cymel®-1170, 100 g (0.206 mole) propyl carbamate, 21.2 g (0.206 mole) acrylic acid, 52.0 g (0.721 mole)	122 g 1.0 propyl 1.5 acrylate	75 °C. 5.5 hrs. 1.5% p-TSA

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PART TWO - PREPARATION OF THE NOVEL VINYL-TERMINATED OLIGOMER OF THE INVENTION (Ingredient B)

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EXAMPLE 18

This Example illustrates the preparation of a vinyl terminated oligomer used in the preparation of compositions of the invention.

A one liter glass reactor was flushed with dry air and charged with the following ingredients:

0.02 grams of phenothiazine

140.6 grams of meta-tetramethylxylylene diisocyanate

The contents of the reaction flask were heated at 80°C. for 10 minutes and 0.15 grams of T-12 (dibutyltin dilaurate) catalyst added. Thereafter, 73.4 grams of 2-hydroxyethyl acrylate were gradually added over a period of thirty minutes and the reaction mixture stirred for one hour at 80 °C.

143 grams of polycaprolactone diol polymer (CAPA 200 diol, molecular weight 550; product of Interox Chemicals Ltd.) were added to the reaction mixture in 4 portions over a period of thirty minutes. Stirring of the reaction mixture was continued for another one and one-half hours at 80°C.

320 grams fo reaction product of urethane acrylates was collected and labeled, "H Oligomer".

EXAMPLE 19

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This Example illustrates the preparation of a vinyl-terminated oligomer used in the preparation of compositions of the invention.

A 500 ml glass reactor was flushed with dry air and charged with the following ingredients:

0.02 grams of benzoquinone

14.6 grams of meta-tetramethyl xylylene diisocyanate

The contents of the reaction flask were heated at 80 °C. for 10 minutes and 0.10 grams of T-12 (Dibutyltin dilaurate) catalyst added. Thereafter, 7.66 grams of 2-hydroxyethyl acrylate were gradually added over a period of thirty minutes and the reaction mixture stirred for one hour at 80 °C.

90 grams of poly(butadiene-acrylonitrile) copolymer (Hycar HTBN 1300x 29, product of Goodrich Tire Company) were added to the reaction mixture in 6 portions over a period of thirty minutes. Stirring of the reaction mixture was continued for another one and one-half hours at 80°C.

100 grams of reaction product of urethane acrylates were collected and labeled, "HT-M Oligomer".

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EXAMPLE 20

A one liter flask was flushed with dry air and charged with 0.06g of benzoquinone and 43.8g of tetramethylxylylene diisocyanate. The mixture was stirred for 10 minutes at 80°C. An 0.3g of T-12 (dibutyltin dilaurate) was added to the reaction mixture, and immediately added 26.6g of hydroxypropyl methacrylate over a period of 30 minutes. The reaction mixture was stirred at 80°C for one hour, then 270g of Hycar HTBN 1300 x 29 (polybutadiene-acrylonitrile) copolymer was added in six portions within 30 minutes. Stirring was continued at 80°C, for another 1 and 1/2 hours. The reaction was thereafter stopped and the reaction products were poured into suitable containers while hot. 330 grams of reaction product of urethane methacrylates were collected and labelled, "HT-MM Oligomer".

PART THREE - TESTS OF THE ADHESIVE PROPERTIES OF THE COMPOSITIONS OF THE INVENTION

SINGLE COMPONENT SYSTEMS

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EXAMPLE 21

Basetock adhesives were prepared from a mixture of 70 parts of HT-M, product of Example 19, 25 parts of M3, product of Example 4, 5 parts of N-vinylpyrrolidone (NVP), 5 parts of succinic anhydride, and 30 parts of IMSIL® 15 Silica, product of Illinois Minerals Co. Several one package formulations comprising the above materials, various initiators, and stabilizers, were used to bound Rockwell 9468 Polyester Plastic Panels SMC. The assemblies were press heated at 250°F. (121.1°C.) for two minutes and the cooled assemblies were postbaked at 110°C./30 minutes. Wedge tests for the basestock adhesives were conducted as follows:

SMC TO SMC BONDING

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The following test procedure was used to evaluate SMC to SMC adhesive bonding for Example 21 and all following Examples.

Commercial polyester plastic SMC panels (From suppliers such as Rockwell International, Diversitech, etc.) were cut into 1" by 4" (2.54 cm by 10.16 cm) coupons for bonding test uses. The bond area of the coupons were wiped with a dry rag without any pretreatment. Enough adhesive was applied to cover a 1" by 1" (2.54 cm by 2.54 cm) overlap area. 30 mil (0.762 mm.) glass beads were scattered on surface of the adhesive (in an amount not to exceed covering more than 1% of surface area). 30 mil spacers (0.762 mm.) (metal shim or Teflon® strip) was used in one end of the test assembly to assure a uniform 30 mil (0.762 mm.) adhesive bond line. The wedge assemblies with spacers inserted and wrapped with a protective mylar film web is made ready for placement in a heated press. The assemblies were press heated 1-2 minutes at 250°F (121.1°C.). Desirable pressure is usually 8-10 psi (6.56 to .70 kg/cm²). After the initial 1-2 minute heating the assemblies were tested for green strength by letting the assembled coupons cool to about room temperature and thereafter hand wiggling the coupons to see that they do not readily separate (pass the test). Thereafter the spacer was removed and excessive adhesive trimmed from exposed edges. Finally, the bonded assemblies were postbaked at 300° F (148.9° C.) for 30 minutes, then removed and cooled. A 45° angle wedge test of the assemblies was performed to determine the type of adhesive failure modes. Types of failure modes were reported as follows: Fiber Tear (FT), Adhesive Failure (AF), Cohesive Failure (CF), Stock Break (SB).

For the lap shear test, a lap configuration of assemblies was used. The adhesive bonds were 1" x 1" (2.54 cm by 2.54 cm.) with 30 mil (0.76 mm.) spacers. The assemblies were press heated at 250° F. (121.1° C.) for 1-2 minutes to observe green strength. All bonded assemblies were postbaked at 300° F. (148.9° C.) for 30 minutes. The assemblies were removed and cooled. A lap shear measurement by Instron machine was performed at a crosshead speed of 0.05 in/min. (1.27 cm/min) by ASTM Method D-1002).

The test results obtained from the single component system of Example 21 are shown in the following table:

•		Composit	tions	Wedge Result
Formulation	Base Adhesive X	Initiator	Stabilizer	
21A 21B 21C	100 part 100 part 100 part	1 part Vazo-52 1 part Vazo-64 1 part Benzoyl peroxide	220 ppm p-benzoquinone (BQ) 10 ppm BQ 50 ppm BQ	95% FT 90% FT 100% FT 95% FT
21D 21E	100 part 100 part	1 part Vazo-52 1 part Vazo-64	450 ppm BQ 200 ppm BQ	100% F

Va26 62 9, Va26 6 1 9, Gatzly 21,

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EXAMPLE 22

The above formulations 21 A to E were held in room temperature storage. The aged adhesives were then checked by the same procedures as above in Example 42. Wedge results of Rockwell 9468 SMC-to-SMC bonding test obtained for formulations 21 A to E are listed below. The results show that the adhesive formulations are stable at room temperature up to 90 days.

Formulation	W	/edge Result	s
,	30 day storage	45 day storage	90 day storage
21A 21B 21C 21D 21E	100% FT 80% FT 90% FT	85% FT S.B. 95% FT 95% FT 85% FT	95% FT 95% FT 90% FT

EXAMPLE 23

The following adhesive formulations with different compositions were used to bond SMC coupons (rag wipe only). The assemblies were press heated at 250° F. (121.1° C.) for one minute. The cooled assemblies were postbaked at 300° F. (148.9° C.) for 30 minutes. The wedge test results are tabulated below:

				1044	NAVE TO	TABLE OF EXAMPLE 31 DESTILLE	11 TC		
				I WOLL	ייין דיין	מו דר כו יובס			
				Colu	Compositions				
Formulation	Ą	89	၁	a	ш	Fillers	Others	Substrate	Wedge Result
23A	M3	HT-M	NVP	V-64	S.A.	Imsil/A-15	p-8Q	Rockwell 9468	100% F.T.
	24.60g	64.36g	12.72g	2.119	4.56g	104.83g	200 ppm	Diversitech 0113	95% F.T.
238	M3	M-TH	NP	٧-64	S.A.	Imsil/A-15	p-80	Rockwell 9468	100% F.1.
	24.60g	64.369	12.72g	2.11g	4.56g	104.83g	200 ppm		!
230	M3	M-TH	STY	٨-64	M.A.	Imsil/A-15	Cab-o-sil® TS720	Rockwell 9468	95% F.1.
	12.41g	30.23g	6.03g	0.98g	2.15g	47.22g	0.989		1
23D	M3	HT-M	NVP	٨-64	S.A.	Talc 156	Cab-o-sil® TS720	Rockwell 9468	95% F.T.
	36.400	96.60g	18.80g	2.75g	6.80g	111.71g	2.75g		,
23E	M3	HT-M	NVP	V-64	S.A.	Imsil/A-15		Rockwell 9468	80% F.T.
!	19.91g	27.339	5.40g	0.89g	1.94g	44.53g			
A, B, C, D, E	as defined	in text, Part	t III - the es	sential in	gredients	of the inventic	A, B, C, D, E as defined in text, Part III - the essential ingredients of the invention for adhesive compositions	sitions	
S.A. = succinic anhydride	iic anhydric	eţ							
M.A. = maleic anhydride	c anhydride	Ф.							
p-BQ = p-benzoquinone	nzoquinone								
NVP = N-vinyl pyrrolidone	yl pyrrolido.	ne							
STY = styrene	Φ								
CAB-O-SIL®, fumed silica, product of Cabot Corp.	fumed silic	a, product	of Cabot C	orp.					
IMSIL®. A silica, product of Illinois Minerals Co.	ca. product	of Illinois !	Minerals Co	Ċ					

EXAMPLE 24

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Formulations 23A and 24B previously described were used to bond Rockwell 9468 SMC coupons. The adhesives were cured in the same fashion as Example 23. Lap shear measurements of the cured assemblies were performed at 180° F. (82.2° C.) The data are listed below.

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Formulation	Lap shear data
23A	> 330 psi ⁻ , (23.20 Kg/cm ²) S.B. 340 psi , (23.90 Kg/cm ²) 95% F.T.
23B	> 360 psi*, (25.31 Kg/cm²) S.B. 320 psi , (22.50 Kg/cm²) 100% F.T.

*The lap shear strength measurement could not be continued because stock broke.

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EXAMPLE 25

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The above formulations 23A and 23B were used to bound Rockwell 9468 SMC coupons. The adhesives were cured in the same fashion as in Example 23. The cured assemblies were soaked in the deionized water at 54 °C. for 7 days. At the end of the seventh day, the assemblies were air dried and subject to wedge test and lap shear measurement at room temperature. The results are tabulated below.

Formulation	Wedge result	Lap shear data
23A	100% F.T.	340 psi, (25.31 Kg/cm²) 100% F.T.
11B	95% F.T.	450 psi, (31.64 Kg/cm²) S.B. (100% F.T.)

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TWO COMPONENT SYSTEMS

EXAMPLES 26 to 38

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Two base stock adhesives, 26A and 26B, were formulated according to the following compositions.

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Formulation				Composi	tions	
26A	HT-M 96.60g	M3 36.40g	NVP 18.80g	S.A. 6.80g	Imsil® A15 155.43g	Cab-o-sil® TS720 3.17g
26B	HT-MM 96.70g	M3 36.10g	NVP 18.80g	S.A. 6.80g	imsil A15 155.23g	Cab-o-sil® TS720 3.17g
HT-M is a vin Example 19.	yl-terminate	d polyureth	nane/polyar	nide oligo	omer prepared	by the method of
M3 is a melar	nine derivat	ive prepare	ed by the m	nethod of	Example 4.	

The two component adhesive system comprises:

Part A with initiators (e.g., hydroperoxides, or peroxides); and

Part B with promoters/accelerators (e.g., transition metal complexes, or tertiary aromatic amines). Several two-component adhesives were formulated by using base adhesives 26A and 26B with different initiator/promoter combinations. An equal weight of Part A and Part B formulations were mixed thoroughly. The well-mixed adhesives were applied to Rockwell 9468 SMC coupons. The cure conditions were the same as Example 23. The resultant pot life of the mixed adhesives and the results of wedge test on the assemblies are tabulated below.

IRT A Part B Pot Liffe IA 10g 26A 10 g >60 min. IPO 0.20 g CONaph 0.40 g >60 min. IPO 0.20 g CONaph 0.40 g >60 min. SA 10 g CONaph 0.40 g >60 min. SA 10 g CONaph 0.40 g >60 min. SB 10 g CONaph 0.40 g >60 min. SB 10 g CONaph 0.40 g >60 min. SB 10 g CONaph 0.30 g 26A 10g, CONaph 0.30 g 90 min. GA 10g, CHPO 0.40 g 26A 10g, CONaph 0.30 g 55 min. GA 10g, CHPO 0.40 g 26A 10g, CONaph 0.40 g 45 min.	TABLE OF RES	TABLE OF RESULTS FOR TWO COMPONENT SYSTEMS		5 .
10 g CONaph 0.40 g >60 10 g CONaph 0.40 g pm 10 g CONaph 0.30 g 26A 10 g CONaph 0.30 g 55 10 g CHPO 0.40 g 26A 10 g CONaph 0.40 g 45	IGA	rt B	Pot Life	Wedge Results
10 g CONaph 0.40 g >60 10 g CONaph 0.40 g	2 67 COb	10 g aph 0.40	>60 min.	100% F.T.
10 g CONaph 0.40 g >60 400 ppm p-BQ 100 ppm 10 g CONaph 0.40 g CONaph 0.40 g CONaph 0.40 g P-BQ 200 ppm 10g, CHPO 0.30 g 26A 10g, CONaph 0.30 g 55 10g, CHPO 0.40 g 26A 10g, CONaph 0.30 g 55 10g, CHPO 0.40 g 26A 10g, CONaph 0.40 g 55		10 9	>60 min.	100% F.T.
10 g CoNaph 0.40 g 600 ppm p-BQ 200 ppm g 26A 10g, CONaph 0.30 g 90 10g, CHPO 0.40 g 26A 10g, CONaph 0.40 g 55 10g, CHPO 0.40 g 26A 10g, CONaph 0.40 g 45		A 10 9 Naph 0.40 9 BQ 100 ppm	>60 min.	95 % P. T.
CHPO 0.30 g 26A 10g, CoNaph 0.30 g 90 CHPO 0.40 g 26A 10g, CONaph 0.30 g 55 CHPO 0.40 g 26A 10g, CoNaph 0.40 g 45		18 10 g Maph 0.40 g -80 200 ppm	>60 min.	99% F.T.
6A 10g, CHPO 0.40 g 26A 10g, CONaph 0.30 g 6A 10g, CHPO 0.40 g 26A 10g, CONaph 0.40 g	5	10g, CoNaph 0.30	90 min.	100% F.T.
6A 10g, CHPO 0.40 g 26A 10g, CoNaph 0.40 g	0.40 g	conaph 0.30	min	100% F.T.
D 04.0 The say and Cobest 0.40 G	0.40 g	CONaph 0.40	45 min.	100% F.T.
d zer rohr corem of the	CHPO 0.40 g 26	26A 10g, CoPent 0.40 g	20 min.	100% F.T.

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Pormulation	Part	«						Part B	m				Pot	Pot Life	Wedge Results	sults
35	26A 10	97	76	g, CHPO 0.40 g	0	9.	5	26A	109,	26A 10g, CoPent 0.20 g	0.2	6 0	3	35 min.	100% F.T.	÷.
36	26B 10	2	6	g, CPO 0.40 g	•	9.	5	26B	26B 10g, DMA	DHA	0.40 9	6 0	9 ^	>60 min.	100% F.T.	H
37	26B 1(91	6	g,t-BPO 0.40 g	0	9	б	26B	26B 10g, DHA	DHA	0.40 9	6 0) V	>60 min.	100% F.T.	F
38	26B 10 Lupers	12	\$ ק	0 g, sol DDM-9 0.08 g	0	.08	5	26B	109,	26B 10g, Conaph 0.14 g	0.1	6 1	9 ^	>60 min.	100% F.T.	E

CHPO = cumyl hydroperoxide

CoNaph = 50% wt Cobalt(II) naphthenate solution in NVP

CoPent = Cobalt(II) pentanedionate

CPO = di-cumyl peroxide

DMA = Dimethyl aniline

t-BPO = di-t-butyl peroxide

o = bensoyl peroxide

GLASS TO STEEL BONDING

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The following procedure was employed to evaluate the adhesives fo the invention for glass to steel bonding applications:

Substrates:

 $1" \times 4" \times 1/4"$ (2.54 cm x 10.16 cm x 1.27 cm) plate glass coupons and $1" \times 4"$ (2.54 cm x 10.16 cm) #25 cold rolled GM steel coupons.

Pretreatment:

Glass coupons were dipped into silane I solution (as described below) for 5 mintes. The air dry coupons were baked at 110 °C. for 12 minutes. Steel coupons were treated with tin hydrosol (as described below) for 5 minutes and dried in open air. Then the steel coupons were dipped into silane I solution for 5 minutes and dried in air, followed by baking at 110 °C. for 12 minutes. The adhesives of various formulations were applied in 1/2" by 1" (1.27 cm by 2.54 cm) overlap with 2 mils (0.051 mm.) of thickness. The adhesives were cured at 110 °C. for 30 minutes.

(1) Silane I solution - was prepared by adding to a 500 ml beaker 139.7 g. of absolute ethanol and 74.9 g. of water, and the pH adjusted with aqueous acetic acid to 5. 5.3 g. of 3-methacryloxypropyl trimethoxysilane was added dropwise into the solution with stirring for 15 minutes. The final solution was used for pretreatment of substrates as described above.

(2) Tin hydrosol - was prepared by dissolving 1 g. of SnCl₄H₂O in deionized water to make one weight percent solution. Stannous chloride was then added to the resultant solution in two steps: (i) 2.5 g. SnCl₂ was added, then (ii) 1.0 g. SnCl₂ was added. Each SnCl₂ addition was carried out at room temperature with stirring to ensure complete dissolution before the next addition. The final tin hydrosol solution was used for pretreatment of substrates as described above. The lap shear data are tabulated in the following Table:

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TABLE 4

	T	EST RESU	JLTS OF E	XAMPLES	39 TO 44	
Example	Adhesive Formulation		Compositions			Lap shear strength
		Α	В	C	D	
39	1	M 0.40 g	H 1.30g	NVP 0.30g	V-52 0.02g	2100 psi (147.6 Kg/cm²)
40	J	M1 0.4 g	H 1.30g	NVP 0.30a	V-52 0.02g	1900 psi (119.5 Kg/cm²)
41	к	M2 0.4 g	H 1.30 g	NVP 0.30 g	V-52 0.02 g	1400 psi (98.4 Kg/cm²)
42	11	M 0.1 g	H 1.30 g	NVP 0.60 g	V-52 0.02 g	2100 psi (147.6 Kg/cm²)
43	J1	M1 0.1 g	H 1.30 g	NVP 0.60 g	V-52 0.02 g	1400 psi (98.4 Kg/cm²)
44	K1	M2 0.1 g	H 1.30 g	0.60 g	V-52 0.02 g	1000 psi (70.3 Kg/cm²)

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The results show the superior performance of carbamylated acryl adhesives containing M (formulations I and I1) and M1 (formulations J and J1) over the non-carbamylated ones containing M2 (formulations K and

K1).

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GLASS TO GLASS BONDING

Substrates:

0 1" x 4" x 1/4" (2.54 cm by 10.16 cm by 0.635 cm) plate glass coupons

Pretreatment:

Glass coupons were dipped into Silane I solution for 5 minutes. The air dried coupons were baked at 110 °C. for 12 minutes. To the cooled coupons were applied adhesive in a 1" by 1/2" (2.54 cm by 1.27 cm) overlap 2 mils (0.051 mm.) thick. The adhesives were cured at 110 °C. for 30 minutes.

The lap shear data are listed below.

Example	Adhesive Formulation	Lap shear strength
45	ı	1900 psi (133.6 Kg/cm²)
46	J	1500 psi (105.5 Kg/cm²)
47	К	400 psi (28.1 Kg/cm²)

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The results showed that adhesives containing crosslinkers M and M1 of this invention gave better bond strength than M2, the non-carbamylated acrymelamine.

STEEL TO STEEL BONDING

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Substrate:

1" by 4" (2.54 cm by 10.16 cm) #25 cold rolled GM steel coupons.

Pretreatment:

Steel coupons were degreased by CH₂Cl₂ wipe, followed by dipping the coupons into silane II solution (as described above) for 5 minutes. The air dry coupons were baked at 110 °C. for 12 minutes. These pretreated coupons were used for application of adhesives. Adhesive I was applied on the above pretreated steel coupons with a 1" x 1/2" (2.54 cm by 1.27 cm) overlap. Adhesive thickness was 5 mil (0.127 mm.). The adhesive-bound coupons were cured at 110 °C. for 30 minutes. Adhesives J and K were applied and cured in a similar fashion.

Silane II solution - was prepared as Silane I solution, supra, except that 139.7 g. of absolute ethanol, 7.4 g. of water, and 3 g. of alpha-aminopropyltriethoxy silane were used. The final solution was used for pretreatment of substrates as described below.

The lap shear data to the cured coupons by Instron testing were tabulated below:

Example	Adhesive Formulation	Lap shear strength
48	1	1600 psi (1) (112.5 Kg/cm²)
49	J	1350 psi (94.9 Kg/cm²)
50	К	1100 psi (77.3 Kg/cm²)

(1)psi = pounds per square inch.

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The results show the bond strength of adhesives containing vinyl-terminated carbamylalkylated melamines M and M1, is stronger than non-carbamyl-alkylated M2.

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Overbake Effect on Adhesive Bonds

Substrate:

1" x 4" (2.54 cm by 10.16 cm) #25 cold rolled steel coupons

Pretreatment:

Following the same procedures as in Examples 48-50 for steel pretreatment, the adhesives are applied in 1/2" x 1" (1.27 cm by 2.54 cm) overlap 2 mils (0.0508 mm.) thick, and cured at 110° C. for 30 minutes. Some cured specimen were further overbaked at 200°C. for another 30 minutes. The lap shear data of both 110° C./30 minutes cure and 200° C./30 minutes overbake samples are summarized below.

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Cure conditions of Adhesive Formulations				
Example	Formulation	100° C./30 min	200 ° C./30 min	
		cure	overbake	
51	ı	2250 psi (158.2 Kg/cm²)	2150 psi (151.2 Kg/cm²)	
52	К	2150 psi (151.2 Kg/cm²)	1800 psi (126.6 Kg/cm²)	

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The results show that the adhesives containing carbamylated acrylate melamine compound of this invention (M in formulation I) retained 95% of original bond strength after overbake at 200 C./30 min. However, the adhesives containing non-carbamylated acrylated melamine (M2 in formulation K) lost 16% of its original bond strength after overbake. These results show better thermal stability of adhesives prepared from the novel ingredients and combination of essential ingredients used in this invention over those prepared from analogous non-carbamylated compounds.

It is to be expected that numerous modifications will readily become apparent to those skilled in the art upon reading this description. All such modifications are intended to be included within the scope of the invention as defined in the appended claims.

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Claims

- 1. A nitrogen containing heterocyclic compound containing at least two vinyl-terminated substituents and at least one carbamylmethyl substituent.
- 2. The nitrogen containing heterocyclic compound of Claim 1 wherein the vinyl-terminated substituents and the carbamylmethyl substituent is attached to a nitrogen containing heterocyclic nucleus selected from the group consisting of melamine, oligomers of melamine, benzoguanamine, oligomers of benzoguanamine, glycoluril, oligomers of glycoluril, and mixtures thereof.

3. The substituted triaminotriazine compound represented by the formula:

wherein the substituent R is as follows:

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R is acrylamido, acrylate, allyloxy, methacrylate, methallyloxy, acryloyloxy, acryloyloxyalkoxy, methacryloyloxy, methacrylamido, or -OR⁶, or carbamyl; wherein R⁶ is hydrogen or a C₁ to C₁₈ aliphatic, alicyclic, or aromatic radical; and provided that at least one R substituent is carbamyl, and that at least two R substituents are selected from acrylamido, acrylate, allyloxy, methacrylate, methallyloxy, acryloyloxy, acryloyloxyalkoxy,

methacryloyloxyalkoxy, methacryloyloxy, and methacrylamido.

4. A melamine-type compound represented by formula: $C_3 N_6 (CH_2)_a (H)_{(6-a)} (R^1)_m (R^2)_n (R^3)_{a\cdot (n+m)}$;

and

and

where a = 3 to 6; wherein R1 is a substituent selected from the group

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R4

CH2=C-C-O-

and

25 R4 CH₂=C-C-O-Z

40 0

R⁴
| CH = C-CNH

CH₂=C-CNH ,

56 R⁴

or mixtures thereof, and wherein Z is CH2CH2-O- or

and wherein R⁴ is a hydrogen or a C₁ to C₁₈ alkyl radical, and wherein R² is a carbamyl radical of the formula:

0 R5-0 C -NH-

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wherein R^5 is a C_1 to C_{18} alkyl, alicyclic, hydroxyalkyl, alkoxyalkyl or aromatic radical, and wherein R^3 is $-OR^6$; wherein R^6 is a hydrogen or C_1 to C_{18} aliphatic, alicyclic, or aromatic radical; with the proviso that m is at least 1, and the sum of (m+n) is at least 3.

5. A compound selected from the group consisting of the following: $N^2,N^4,N^6-tris(acrylamidomethyl)-N^2,N^4,N^6-tris(2-ethylhexylcarbamylmethyl) melamine$ N2,N4,N5-tris(2-ethylhexylcarbamylmethyl)--N2,N4,N5-tris(2-acryloyloxyethoxymethyl)melamine N2,N4-bis(methylcarbamylmethyl)-N2,N4,N6,N6-tetrakis(2-acryloyloxyethoxymethyl)melamine N²,N⁴-bis(methylcarbamylmethyl)-N²,N⁴,N⁶,N⁶-tetrakis(2-methacryloyloxyethoxymethyl)melamine N2,N4,N5-tris(propylcarbamylmethyl)-N2,N4,N5-tris(2-acryloyloxyethoxymethyl)melamine N2,N4,bis(propylcarbamylmethyl)-N2,N4,N6,N6-tetrakis(2-acryloyloxyethoxymethyl)melamine N2,N4,N6-tris(propylcarbamylmethyl)-N2,N4,N6-tris(2-methacryloyloxyethoxymethyl)melamine N2,N4,bis(methylcarbamylmethyl)-N2,N4,N6,N6-tetrakis(2-methacryloyloxypropyloxymethyl)melamine N², N⁴, N⁶-tris(acrylamidomethyl)-N², N⁴, N⁶-tris(methylcarbamylmethyl)melamine N².N⁴.N⁵-tris(methylcarbamylmethyl)-N²,N⁴,N⁶-tris(2-acryloloxyethoxymethyl)melamine N^2,N^4,N^6 -tris(methylcarbamylmethyl)- N^2,N^4,N^6 -tris(2-methacryloloxyethoxymethyl)melamine $N^2, N^4, N^6 - tris(methylcarbamylmethyl) - N^2, N^4, N^6 - tris(2-acryloloxypropyloxymethyl) melamine. \\$ N²,N⁴,N⁶-tris(propylcarbamylmethyl)-N²,N⁴,N⁶-tris(2-methacryloloxypropyloxymethyl)melamine $N^2, N^4 - bis(propylcarbamylmethyl) - N^2, N^4 - N^5 - tetrakis(2-methacryloloxypropyloxymethyl) melamine and the state of the state$ $N^2, N^4 - bis(propylcarbamylmethyl) - N^2, N^4, N^6 - N^6 - tetrakis(2-acryloloxyethoxymethyl) melamine and the state of the state o$ $N^2, N^4, N^6 - tris (propylcarbamylmethyl) - N^2, N^4, N^6 - tris (acrylamidomethyl) melamine, acrylamidomethyl) melamine, acrylamidomethyl (propylcarbamylmethyl) - N^2, N^4, N^6 - tris (acrylamidomethyl) - N^2, N^4, N^6 - tris (acrylami$ and mixtures thereof.

6. A benzoguanamine compound represented by the formula: $(C_6H_5)C_3N_5 - (CH_2)_b(H)_{4-b} - (R^1)_m(R^2)_n(R^3)_{b-(n+m)}$ wherein b = 3 or 4; and wherein R¹ is a substituent selected from the group

R⁴ | CH₂=C-C-O-

and

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CH₂-C-O-Z

and

75 R4 CH₂=C-CNH

and

20 R4 CH2=C-CH2-O- .

or mixtures thereof, and wherein Z is CH2CH2-O- or

-CH₂-CH-CH₃, -CH₂-CH₂, -CH-CH₂-CH₃,

and wherein R^4 is a hydrogen or a C_1 to C_{18} alkyl radical, and wherein R^2 is a carbamyl radical of the formula:

0 R5-0 **C** -NH-

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wherein R^5 is a C_1 to C_{18} alkyl, alicyclic, hydroxyalkyl, alkoxyalkyl or aromatic radical, and wherein R^3 is -OR⁶; wherein R^6 is a hydrogen or C_1 to C_{18} aliphatic, alicyclic, or aromatic radical; with the proviso that m is at least 2, n is at least 1, and the sum of (m+n) is at least 3.

7. A glycoluril type compound represented by the formula:

 $C_4 H_2 N_4(O)_2$ - $(CH_2)_c(H)_{4-c}$ - $(R^1)_m(R^2)_n(R^3)_{c-(m+n)}$

wherein C is 3 to 4; and wherein R1 is a substituent selected from the group

R⁴
|
CH₂=C-C-O-

and

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R4 | |CH₂=C-C-O-Z

and

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R⁴

|
CH₂=C-CNH

"
O

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and

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R⁴
|
CH₂=C-CH₂-O-

or mixtures thereof, and wherein Z is CH2CH2-O- or

-CH₂-CH-CH₃

-CH₂-CH₂-CH₂

-сн-сн₂-сн₃ | | | |

and wherein R^4 is a hydrogen or a C_1 to C_{18} alkyl radical, and wherein R^2 is a carbamyl radical of the formula:

5 R5-O C -NH-

wherein R^5 is a C_1 to C_{18} alkyl, alicyclic, hydroxyalkyl, alkoxyalkyl or aromatic radical, and wherein R^3 is $-OR^6$; wherein R^6 is a hydrogen or C_1 to C_{18} aliphatic, alicyclic, or aromatic radical; with the proviso that m is at least 2, n is at least 1, and the sum of (m+n) is at least 3.

- 8. A vinyl-terminated polyurethane/polyamide polymer having a molecular weight of from about 3000 to about 80000, containing at least two vinyl end groups; said polymer produced by the process of reacting polyisocyanate with a vinyl acid, vinyl alcohol, or vinyl amide; then, in a second step reacting the first step reaction product with a high molecular weight polyol.
 - 9. A curable coating/sealant composition comprising:
- A. from about 5 to about 70 weight percent of at least one nitrogen containing heterocyclic compound having as reactive groups at least two vinyl-terminated organic groups, and at least one carbamylmethyl group:
- B. from about 30 to about 90 weight percent of a vinyl-terminated polyurethane/polyamide polymer having a molecular weight from about 3000 to about 80,000;
 - C. from about 2 to about 30 weight percent of a polymerizable diluent;
 - D. from about 0.5 to about 5 weight percent of a free-radical catalyst;
 - wherein the sum of A, B, C and D does not exceed 100 weight percent.
 - 10. A curable adhesive composition comprising:
- 5 A. from about 5 to about 70 weight percent of at least one nitrogen compound having as reactive groups at least two vinyl-terminated organic groups, and at least one carbamylmethyl group;
 - B. from about 30 to about 90 weight percent of a vinyl-terminated polyurethane/polyamide polymer having a molecular weight from about 3000 to about 80,000;
 - C. from about 2 to about 30 weight percent of a polymerizable diluent;
- D. from about 0.5 to about 5 weight percent of a free-radical catalyst;
 - E. from about 0.5 to about 10 weight percent of an adhesion promoter/coupling agent; wherein the sum of
 - A, B, C, D and E does not exceed 100 weight percent.

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- Novel vinyl-terminated carbamylmethylated melamines, vinyl-terminated polyurethane/polyamide polymers and coating/adhesive compositions containing same.

⑤ Disclosed are novel nitrogen containing heterocyclic compounds and novel vinyl terminated polyurethane/polyamide polymers together with the use of both of these compounds in a novel adhesive and coating composition. The nitrogen containing heterocyclic compounds of the invention contain two vinyl-terminated substituents and at least one carbamylmethyl substituent and have a nucleus selected from melamine, oligomers of melamine, benzoguanamine, and oligomers of benzoguanamine, glycoluril and oligomers of glycoluril. The vinyl terminated polyurethane compounds of the invention have a molecular weight of from about 3000 to about 80000, and contain at least two vinyl end groups.



EUROPEAN SEARCH REPORT

EP 89 11 4816

	OCUMENTS CONSID	ndication, where appropriate,	Relevant	CLASSIFICATION OF THE
ategory		ndication, where appropriate, it passages	to claim	APPLICATION (Int. Cl.5)
A	EP-A-0 006 206 (TEIJIN LTD claims 1, 3, 5 *).)	1-4,9	C 07 D 251/54 C 07 D 251/18 C 07 D 487/04
A,D	US-A-4 230 550 (K.D. VINCE column 2, lines 20 - 36 * * ab		1-4,9	C 08 G 18/67 C 08 L 75/16
A,D	EP-A-0 245 700 (AMERICAN * claims 1, 5 & US-A-4710542	•	1,4,6,9	C 09 D 175/16 C 09 J 175/16 C 08 K 5/3492
A,D	EP-A-0 245 701 (AMERICAN * claims 1, 5 & US-A-4708984	· · · · · · · · · · · · · · · · · · ·	1,4,6,9	C 08 K 5/3445// (C 07 D 487/04 C 07 D
Α	US-A-2 694 687 (P.L. DE BE * column 1, lines 42 - 49; clair -	•	1-3,9	C 07 D 235:00 C 07 D 235:00)
Α	EP-A-0 044 150 (WESTING) * claims 1, 8 *	HOUSE ELECTRIC CORP.)	1,2,9	
A,D	US-A-4 230 740 (J.D. MOYE * abstract *	ER) 	1-4	TECHNICAL FIELDS
Α	US-A-3 419 535 (P. SCHLU * claim 7 *	MBOM) 	1-4	SEARCHED (Int. Cl.5) C 07 D 251/00
P,X	EP-A-0 314 489 (HOECHST relaim 1; page 5, example 4		8	C 08 F 299/00 C 08 G 18/00
х	EP-A-0 076 126 (OLIN COF * claims 1, 6, 11, 12 *	P.)	. 8	
Y	EP-A-0 228 854 (INTEREZ, * claims 1, 2 *	INC.)	8	
		-/-		
	The present search report has be	en drawn up for all claims		
-	Place of search	Date of completion of search	- '	Examiner
1	Berlin	14 May 91		HASS C V F

- Y: particularly relevant it combined with another document of the same catagory
 A: technological background
 O: non-written disclosure
 P: Intermediate document
 T: theory or principle underlying the invention

- L: document cited for other reasons
- &: member of the same patent family, corresponding document



EUROPEAN SEARCH REPORT

Application Number

EP 89 11 4816

gory		Indication, where appropriate, ant passages	Rele to c	vant laim	CLASSIFICATION OF THE APPLICATION (Int. CL5)
,	EP-A-0 145 929 (DESOTO, * claims 1, 4 *	INC.)	8		
	GB-A-2 010 875 (AMERICA * claims 1, 11 *	AN CYANAMID CO.)	9		
	DE-A-2 009 621 (BORDEN claims 1, 3 *	, INC.)	10		
		· 			
				_	TECHNICAL FIELDS
				-	SEARCHED (Int. Cl.5)
	The present search report has	been drawn up for all claims Date of completion of	search		Examiner
	Berlin	14 May 91			HASS C V F
Υ:	CATEGORY OF CITED DOC : particularly relevant if taken alone : particularly relevant if combined wi document of the same category		the filing di D: document d L: document	ate cited in the cited for otl	
0	: technological background : non-written disclosure : intermediate document				eatent family, corresponding

